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CLAIMS

Peptides, or fragments thereof, as stimulators of insulin secretion and pancreatic beta cell function, the, or each, peptide having at least 50%, preferably at least 80%, more preferably at least 90%, optionally more than 95%, sequence identity based on the ClustalW alignment method with the respective amino acid sequences of SEQ ID Nos. 1 to 17, the peptides having the following structures:

- (a) RRKPLFPFIPRPK (peptide 1.10, Agalychnis calcarifer),
- (f) GKPFYPPPIYPEDM (peptide 24, Bombina variegata),
- (g) IYNAICPCKHCNKCKPGLLAN (peptide 25, Bombina variegata), a peptide having the partial structure ---G-QWA-GH-M, the peptide being preferably selected from (d) Pyr-QRLGHQWAVGHLM-amidated (peptide 21, Bombina variegata) and (e) Pyr-DSFGNQWARGHFM-amidated (peptide 22, Bombina variegata),
- (j) ALSILRGLEKLAKMGIALTNCKATKKC (peptide 3.8, Rana palustris), a peptide having the partial structure FLP--AG-AA---PKIFC-I--KC, the peptide being preferably selected from (k) FLPHAGVAAKVFPKIFCAISKKC (peptide 4.1, Rana pipiens) and (q) FLPLLAGLAANFLPKIFCKITRKC (peptide 8.3, Rana saharica), a peptide having the partial structure A-WKD-LKN-GKAAGKAVLN-VTDMVN-, the peptide being preferably selected from (c)

AVWKDFLKNIGKAAGKAVLNSVTDMVNE (peptide 2.9, Agalychnis litodryas) and (i) ALWKDILKNVGKAAGKAVLNTVTDMVNQ (peptide 2.10, Phyllomedusa trinitatis),

a peptide having the partial structure KG—LL—ASCKLS—C, the partial structure being preferably GIL—LK-FA—AGKG—LL—ASCKLSGQC, the peptide being more preferably selected from (I) KGAAKGLLEVASCKLSKSC (peptide 4.22, Rana saharica), (o) GILSTIKDFAIKAGKGAAKGLLEMASCKLSGQC (peptide 5.6, Rana saharica), (p) GILLDKLKNFAKTAGKGVLQSLLNTASCKLSGQC (peptide 6.5, Rana saharica),

a peptide having the partial structure GIFSK---KK-

KNLLISGLKNVGKEVGMDVVRTGIDIAGCKIKGEC, the peptide being more preferably selected from (m)

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GIFSKFGRKKIKNLLISGLKNVGKEVGMDVVRTGIDIAGCKIKGEC

(peptide 5.1 Rana saharica) and (n)

GIFSKLAGKKLKNLLISGLKNVGKEVGMDVVRTGIDIAGCKIKGEC (peptide 5.4 Rana saharica).

- (b) a peptide having the N-terminus sequence MLADVFEKIMGD... (N-terminus of peptide 1.7, Agalychnis litodryas) and
- (h) a peptide having the N-terminus sequence **XXPLAPFFQAVFK**... (N-terminus of peptide 1.8, *Phyllomedusa trinitatis*).
- Peptides selected from the group comprising brevinins, dermaseptins and esculentins for stimulating insulin secretion by activation of physiological stimulussecretion coupling pathways, rather than by antimicrobial action involving cell lysis.
- 3. A peptide as claimed in claim 1 or 2 with at least one amino acid modification by insertion of fatty acid at the alpha amino group of native amino acid or an epsilon amino group of a substituted lysine residue.
- 4. A peptide as claimed in any one of claims 1, 2 or 3, having at least one amino acid substitution and/or modification including N-glycated, N-alkylated, N-acetylated, N-acylated, N-isopropyl, and / or N-pyroglutamyl amino acids.
- 5. Use of at least one peptide as claimed in any one of claims 1 to 4 in the preparation of a medicament to stimulate insulin secretion and / or moderate blood glucose excursions.
- 6. The use of at least one peptide as claimed in any one of claims 1 to 4 in the preparation of a medicament for treatment of type 1 or type 2 diabetes mellitus.
- 7. A pharmaceutical composition including at least one peptide according to any one of claims 1 to 4 in admixture with a pharmaceutically acceptable excipient.
- 8. A pharmaceutical composition useful in the treatment of obesity and/or type 2 diabetes which comprises an effective amount of at least one peptide as claimed in any of claims 1 to 4 in admixture with a pharmaceutically acceptable excipient, the

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pharmaceutical composition being preferably for delivery through transdermal, nasal inhalation, oral or injected routes.

9. A pharmaceutical composition as claimed in claim 8 which further comprises at least one further pharmaceutically active agent, the, or each, further pharmaceutically active agent being preferably selected from one or more sulphonylureas, meglitinides, metformin, and/or thiazolidinediones, or a mixture thereof.